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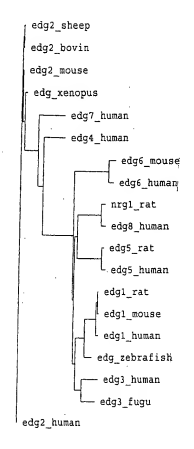
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## (54) EDG8 receptor, its preparation and use

(57) The present invention relates to newly identified EDG 8 receptors, polynucleotides encoding this receptor, polypeptides encoded by such polynucleotides the preparation and the use of thereof.

Fig. 1B



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#### Description

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[0001] The present invention relates to newly identified EGD8 receptors, polynucleotides encoding this receptor, polypeptides encoded by such polynucleotides the preparation and the use of thereof.

[0002] In an effort to identify new G-protein coupled receptors of the EDG (endothelial differentiation gene)-family a novel member of the EDG-family of G-protein coupled receptors, Human EDG8, was identified. The full-length clone was isolated and studies on chromosomal mapping, tissue expression and identification as a functional cellular receptor for sphingosine 1-phosphatea were performed. Taken together, the data provide compelling evidence that EDG8 is the fifth receptor for sphingosine 1-phosphate, exclusively expressed in peripheral tissues, its presence in endothelial cells being responsible for the broad tissue distribution.

[0003] The lysolipid phosphate mediators lysophosphatidic acid (LPA) and sphingosin 1-phosphate (S1P) have attracted increasing attention as modulators of a variety of important biological functions (Moolenaar *et al.*, 1997; Morris, 1999; Lynch, 1999) and their list of biological activities is continuously growing.

Among the biological responses to LPA is platelet aggregation (Durieux and Lynch, 1993; Moolenaar, 1994; Jalink et al., 1994; Siess et al., 1999; Gueguen et al., 1999), smooth muscle contraction, in vivo vasoactive effects, chemotaxis, expression of adhesion molecules on endothelial cells, increased tight junction permeability, activation of membrane ion channels and many others. The biochemical signalling events that mediate the cellular effects of LPA include stimulation of phospholipases, mobilization of intracellular Ca2+, inhibition of adenylyl cyclase, activation of phosphatidylinositol 3-kinase, activation of the Ras-Raf-MAP kinase cascade and stimulation of Rho-GTPases.(Moolenaar et al., 1997)

S1P, in particular, is implicated in cell proliferation, induction/suppression of apoptosis, modulation of cell motility, angiogenesis, tumor invasiveness, platelet activation and neurite retraction. Cellular signalling by S1P involves activation of PLCβ and subsequent intracellular Ca<sup>2+</sup> release, activation of MAP-kinases, activation of inward rectifying K+-channels and inhibition and/or activation of adenylyl cyclase.

Both, LPA and S1P are recognized to signal cells through a set of G-protein coupled receptors (GPCRs) known as EDG (endothelial differentiation gene)-receptors. The EDG-family of GPCRs currently comprises seven human members (EDG1-7) that fall into two major groups depending on their preference for the activating lipid-ligand: EDG1, 3, 5 and 6 preferentially interact with S1P, EDG2, 4 and 7 preferentially interact with LPA.

The assignment of specific biological functions to certain receptor subtypes is hampered by the fact that EDG receptors are expressed in an overlapping fashion, they activate multiple and in part identical signal transduction pathways, the selectivity for their activating ligands is not absolute, and medicinal chemistry is only poorly developed in that specific antagonists for dissecting the pharmacology of the individual subtypes are not available yet.

An important step to shed more light on the biological role of the individual receptor subtypes would be to identify the complete set of receptors that respond to the phospholipid mediators S1P and LPA.

[0004] The present invention relates to newly identified EGD8 receptors, polynucleotides encoding this receptor, polypeptides encoded by such polynucleotides the preparation and the use of thereof.

**[0005]** The present invention relates to an isolated polynucleotide comprising a nucleotide sequence that has at least 80 % identity to a nucleotide sequence encoding the polypeptide of SEQ ID NO. 2 or the corresponding fragment thereof; or a nucleotide sequence complementary to said nucleotide sequence.

[0006] Preferably, the polynucleotide is DNA or RNA.

Preferably, the nucleotide sequence of the polynucleotide is at least 80 % identical to that contained in SEQ ID NO. 1. In another embodiment, the polynucleotide has the nucleotide sequence SEQ ID NO. 1. In another embodiment, the polynucleotide encodes the polypeptide of SEQ ID NO. 2 or a fragment thereof.

[0007] Another aspect to the invention relates to an expression system for the expression of EDG8. The EDG8 DNA or RNA molecule comprising an expression system wherein said expression system is capable of producing a polypeptide or a fragment thereof having at least 80 % identity with a nucleotide sequence encoding the polypeptide of SEQ ID NO. 2 or said fragment when said expression system is present in a compatible host cell.

The invention relates to a host cell comprising the expression system.

[0008] In another aspect, the invention relates to a process for producing a EDG8 polypeptide or fragment comprising culturing a host cell comprising the expression system under conditions sufficient for the production of said polypeptide or fragment. Preferably, the said polypeptide or fragment is expressed at the surface of said cell. The invention relates also to cells produced by this process.

The process preferably further includes recovering the polypeptide or fragment from the culture.

[0009] In another aspect, the invention relates to a process for producing a cell which produces a EDG8 polypeptide or a fragment thereof comprising transforming or transfecting a host cell with the expression system such that the host cell, under appropriate culture conditions, produces a EDG8 polypeptide or fragment.

[0010] In particular, the invention relates to an EDG8 polypeptide or a fragment thereof comprising an amino acid sequence which is at least 80 % identical to the amino acid sequence contained in SEQ ID NO. 2; in particular to an

EDG8 polypeptide or a fragment thereof having amino acid sequence SEQ ID NO. 2 or a part thereof.

[0011] Further, the invention relates to a process for diagnosing a disease or a susceptibility to a disease related to expression or acitivity of EDG8 polypeptide comprising:

- a) determining the presence or absence of mutation in the nucleotide sequence encoding said EDG8 polypeptide in the genome of said subject; and/or
- b) analyzing for the presence or amount of the EDG8 polypeptide expression in a sample derived from said subject.

[0012] In addition, the invention relates to a method for identifying compounds which bind to EDG8 polypeptide comprising:

- a) contacting a cell comprising the expression system or a part of such a cell with a candidate compound; and
- b) assessing the ability of said candidate compound to bind to said cells.
- [0013] Preferably, the method further includes determining whether the candidate compound effects a signal generated by activation of the EDG8 polypeptide at the surface of the cell, wherein a candidate compound which effects production of said signal is identified as an agonist.
  - [0014] In another embodiment of the invention, the method further includes determining whether the candidate compound effects a signal generated by activation of the EDG8 polypeptide at the surface of the cell, wherein a candidate compound which effects production of said signal is identified as an antagonist.
  - [0015] The invention also relates to an agonist or antagonist identified by such methods.
  - [0016] In another special embodiment of the method, the method further includes contacting said cell with a known agonist for said EDG8 polypeptide; and determining whether the signal generated by said agonist is diminished in the presence of said candidate compound, wherein a candidate compound which effects a diminution in said signal is identified as an antagonist for said EDG8 polypeptide. The known agonist is for example S1P, LPA and/or DHS1P. The invention also relates to an antagonist identified by the method.
  - [0017] The invention in addition, relates to a method of preparing a pharmaceutical composition comprising
    - a) identifying a compound which is an agonist or an antagonist of EDG8,
    - b) preparing the compound, and

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- c) optionally mixing the compound with suitable additives.
- [0018] The invention also relates to a pharmaceutical compound prepared by such a process.
- [0019] In the study, we report about the cloning, chromosomal mapping, tissue expression and functional identification as a receptor for S1P of a novel GPCR, EDG8, the fifth functional receptor for sphingosine 1-phosphate.
- [0020] In an effort to identify new G-protein coupled receptors of the EDG-family a database search with alignments of the currently known 18 members of this receptor family, comprising human EDG1-7 sequences up to the putative EDGs from Xenopus and Zebra-fish was performed. A multiple alignment of these sequences was created by CLUS-TALW and used in a PSI-BLAST search to scan translated versions of human genomic DNA sequences, which were publicly available in the different EMBL sections. For translation of DNA into protein sequences, individual protein files within two respective STOP-codons were created and all proteins shorter than 50 amino acids were ignored. As the majority of GPCRs is unspliced searching for GPCRs within genomic sequences should bring about novel receptor proteins.
- Performing a PSI-BLAST search, the various cDNAs and genomic contigs, respectively, for the human EDG1-7 receptors were identified, and an additional genomic hit with a high e-value, that was not identical to any of the published EDG-sequences. The nucleotide and amino acid sequence of the new putative GPCR are depicted in Fig.1A. Hydropathy analysis (hydrophobicity plot not shown) suggests a seven transmembrane protein with three alternating extra-and intracellular loops, assumed to be the heptahelix structure common to GPCRs.
- To shed more light on the relationships involved in the molecular evolution of the EDG-receptor family, a grow tree phylogram was constructed using the neighbor joining method (GCG software) (Fig.1B) According to this phylogenetic tree, the human EDG-family can be divided into two distinct groups: EDG1, 3, 5 and 6 belonging to one, EDG2, 4 and 7 belonging to the other group. These two groups are discriminated further by their preference for different lipid ligands: EDG1, 3, 5, 6 are preferentially stimulated by sphingosin 1-phosphate (S1P), EDG2, 4 and 7 by lysophosphatidic acid (LPA). The newly identified sequence was named EDG8 for reasons of consistency with the existing human EDG-family nomenclature and exhibited highest similarity (86.8%) to the rat nrg1-protein (Fig. 1B), a GPCR recently cloned by EST-expression profiling from a rat PC12 cell library (Glickman et al., 1999). The high similarity between EDG8 and the known sphingosin 1-phosphate (S1P) receptors EDG1, 3 and 5 (48-51%) (Fig. 1C) led us to test the hypothesis that EDG8 may be a functional S1P-receptor.

In testing for S1P receptor activity, the EDG8 cDNA was introduced into chinese hamster ovary (CHO) cells by transient transfection. CHO cells were chosen as they exhibit minimal responses to sphingosin 1-phosphate in concentrations up to 1  $\mu$ M but respond to S1P after transfection with the S1P preferring receptors EDG 1, 3 and 5 (Okamoto et al., 1998; Kon et al., 1999). To test functional receptor activity, it was decided to monitor the mobilization of intracellular Ca2+ for three reasons:

1.) S1P has been reported to increase Ca2+ in many cell types;

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- 2.) the identification of EDG1, 3, 5 and 6 as receptors for S1P has provided the molecular basis for a GPCR mediated mechanism and the receptors are known to mediate intracellular Ca2+ through either PTX-sensitive Gi proteins or the PTX-insensitive Gag/11 pathway;
- 3.) all currently known S1P-responding EDG-receptors (except EDG6) are present in endothelial cells (A. Niedernberg et al., submitted), in which intracellular Ca2+ release is a major pathway in the generation of NO, an important factor in vascular biology.

Thus, identification of the complete set of S1P receptors, involved in intracellular Ca2+ mobilization could help clarify the role of the individual subtypes in endothelial cell signalling.

[0021] Fig.2 depicts measurement of the intracellular Ca2+ concentration, mediated by S1P via the different S1Preceptors EDG1, 3, 5, 6 and the putative S1P-receptor EDG8, cotransfected in CHO cells with empty vector DNA as a control or various G-protein α subunits. Intracellular Ca2+ concentrations were recorded as real time measurements using the Fluorescence plate imaging reader (FLIPR, Molecular Devices). As has been reported for EDG1, 3 and 5, S1P elicited intracellular Ca2+ signals that did not require cotransfection of a G-protein α subunit. As already known for a large number of Gq-coupled receptors, coexpression of Gaq augments the EDG1 and 5-mediated Ca2+-release as compared with the Ca2+ signal induced by stimulation of endogeneous Gaq. In case of EDG3, additional exogeneously added Gaq did not further improve the signal intensity. In case of EDG6, Yamazaki Y. et al. (2000) reported an S1P-induced mobilization of intracellular Ca2+ but we failed to detect a significant Ca2+ increase above basal levels in the absence of any cotransfected G-protein α subunit. The reason for this discrepancy could be the cellular background, as Yamazaki Y. et al. (2000) reported that the Ca2+ signal can be completely abolished in the presence of pertussis toxin (PTX), indicating the involvement of Gi-type G-proteins. In this case the Ca2+ signal would be elicited by bg, released from activated Gaibg heterotrimers. The Gαi-induced Ca2+ signals are known to be much smaller in intensity as compared with the Ca2+ signals induced by bona-fide Gq-linked receptors (Kostenis et al., 1997a; Kostenis et al., 1997b). It may be that detection of such Ca2+ concentrations is beyond the sensitivity of the FLIPR system. EDG8 did not release intracellular Ca2+ when stimulated with S1P (Fig.2), but gained the ability to mobilize Ca2+ upon cotransfection with Gα16, a G-protein α-subunit, known to couple GPCRs from different functional classes to the Gq-PLC $\beta$  pathway or G $\alpha$ qi5, a mutant G-protein  $\alpha$  subunit that confers onto Gi-liked receptors the ability to stimulate Gq. These results show that EDG8 as opposed to EDG 3 and 5 is not a bona-fide Gq-coupled receptor but a fuctional cellular receptor for S1P. To check, whether the EDG8 receptor also reacts to related lysophospholipid mediators, we examined the abilities of lysophosphatidic acid (LPA), dihydrosphingosin 1-phosphate (DHS1P), sphingosylphosphorylcholine (SPC) and lysophosphatidylcholine (LPC) to increase intracellular Ca2+ in CHO cells transiently transfected with the EDG8 receptor and the G-protein α subunits G16 and Gqi5 (Fig.3). Besides S1P, which was the most potent activator of EDG8, LPA and DHS1P evoked [Ca2+]i increases in concentrations of 100 and 1000 nM. SPC and LPC, respectively, failed to generate any significant response in concentrations up to 1 μM.. These data show that EDG8 is a S1P preferring receptor, but also responds to related phospholipids like DHS1P or LPA, as has also been reported for EDG1, which is a high affinity receptor for S1P and a low affinity receptor for LPA.

[0022] Next, the expression pattern of the human EDG8 gene in human tissues was investigated by Northern blot analysis (Fig.4). Northern blot analysis shows EDG8 expression in several peripheral tissues. Tissues positive for EDG8 RNA were skeletal muscle, heart and kidney, lower abundance of RNA was seen in liver and placenta, no signal was obtained in brain, thymus, spleen, lung and peripheral blood leukocytes. In all tissues a single RNA transcript of 5.5 kb was observed after hybridization with a DIG-labelled EDG8 antisense RNA probe. EDG8 exhibits highest similarity to the rat nrg1-GPCR with an amino acid identity of 86.8% (Fig.1B) suggesting that it may be the human homolog of the rat nrg1 protein. However, the expression pattern of human EDG8 is quite different from the rat nrg1-receptor, which is found almost exclusively in brain. This finding suggests that EDG8 may represent a closely related but entirely different receptor from nrg1, rather than the human homolog. Never the less, it does not rule out the possibility that EDG8 and nrg1 are homologs with entirely different, species-dependent expression patterns.

[0023] As the first member of the EDG-family of GPCRs was cloned as an immediate early gene induced during the morphogenetic differentiation phase of angiogenesis (Hla and Maciag, 1990) and subsequently cloned from a human umbilical vein endothelial cell library exposed to fluid shear stress as an upregulated gene it is reasonable to assume that EDG receptors play an important role in the regulation of endothelial function. Therefore, the presence of EDG8 transcripts in several human endothelial cell lines was analyzed. RT-PCR analysis of human umbilival vein endothelial

cells (HUVECs), human coronary artery endothelial cells (HCAECs), human microvascular endothelial cells of the lung (HMVEC-L) and human pulmonary artery endothelial cells (HPAEC) revealed EDG8 expression in all cell lines tested (Fig.5A). In Fig.5B it is shown that EDG8 specific primers indeed solely amplify EDG8 sequences and none of the related EDG1-7 sequences. These findings suggest that the presence of EDG8 in different peripheral organs may be due to its localization in endothelial cells; it does not rule out, however, that EDG8 transcripts occur in cell types other than endothelial cells.

[0024] The expression of EDG8 in addition to EDG1, 3, and 5 (Rizza et al., 1999) in HUVECS and several other endothelial cell lines is intriguing in view of all the reports regarding S1P effects on endothelial cell signalling. Hisano et al. (1999) reported that S1P protects HUVECS from apoptosis induced by withdrawal of growth factors and stimulates HUVEC DNA synthesis; the authors derived a model for cell-cell interactions between endothelial cells and platelets but the S1P-receptor responsible for HUVEC-protection of apoptosis could not be identified. Rizza et al., 1999 reported that S1P plays a role in endothelial cell leukocyte interaction in that S1P induces expression of cell adhesion molecules in human aortic endothelial cells, allowing monocytes and neutrophils to attach. These effects were blocked by pertussis toxin, suggesting the involvement of a Gi-coupled S1P receptor. The responsible S1P-receptor subtype, however, could not be identified and the EDG8 receptor was not included at the time of this study. Expression profiling of all EDG receptors in individual cell lines and the use of EDG receptor subtype selective compounds will clearly be necessary to help determine the role of the individual S1P receptors in endothelial cell signalling mechanisms.

[0025] Finally, the mapping of EDG receptors in genomic sequences allowed to allocate a map position for four genes of this family (Tab.1). Interestingly, so far, four EDG-receptors including EDG8 are located on chromosome 19.

In conclusion, we isolated a new member of the EDG-family of G-protein coupled receptors, EDG8, and showed that it functions as a cellular receptor for sphingosine 1-phosphate. EDG8 could exclusively be detected in peripheral tissues like skeletal muscle, heart and kidney and several human endothelial cell lines. It is conceivable that the expression in endothelial cells may account for the broad tissue distribution of this receptor. The existence of at least eight EDG-receptors for lysophospholipids suggests that receptor subtype selective agonists and antagonists will essentially be necessary for a better understanding of the biology of lysophospholipids and their respective receptors.

#### Figure legends

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[0026] Fig.1A: The nucleotide and deduced amino acid sequence of human EDG8. The deduced amino acid sequence is shown below the nucleotide sequence with the nucleotide positions indicated on the left. The putative seven transmembrane domains are underlined.

[0027] Fig. 1B: Phylogenetic tree of the EDG-family of receptors. The phylogenetic tree depicted was derived by the neighbor joining method method performed with the GCG program.

[0028] Fig. 1C: Alignment of the amino acid sequence of human EDG8 with the other EDG-family members. The amino acid sequence of EDG8 is compared with the EDG1-7 polypeptides (EDG1: accession number M 31210, EDG2: accession number U 80811, EDG3: accession number X 83864, EDG4: accession number af 011466, EDGS: accession number af 034780, EDG7: accession number af 127138). The approximate boundaries of the seven putative transmembrane domains are boxed. Gaps are introduced to optimize the alignment.

[0029] Fig.2: Mobilization of intracellular Ca<sup>2+</sup> by S1P mediated by the EDG1, 3, 5 and 8 receptor in CHO cells, cotransfected with empty vector DNA as a control or the indicated G-protein α subunits. Agonist-mediated changes of intracellular Ca<sup>2+</sup> were measured with the FLIPR using the Ca<sup>2+</sup>-sensitive dye FLUO4. Fluorescence of transfected cells loaded with FLUO4 was recorded before and after addition of S1P, applied in the indicated concentrations. Data are expressed as means of quadruplicate determinations in a single experiment. An additional experiment gave similar results

[0030] Fig.3: Effects of S1P, LPA and related lysophospholipid mediators on EDG8-mediated increase in intracellular Ca<sup>2+</sup> measured with the FLIPR as described in the legend of Fig.4. The different lipids were applied in concentrations of 10, 100 and 1000 nM, respectively. Data are means of quadruplicate determinations of a representative experiment. Two additional experiments gave similar results.

[0031] Fig.4: Northern blot analysis of EDG8 in human tissues. Poly(A)+ RNA (1µg) from various human tissues (human multiple tissue Northern blots, CLONTECH) was hybridized with probes specific to human EDG8 (upper panel) and glyceraldehyd-3-phosphate dehydrogenase, GAPDH, (lower panel) on a nylon membrane. The origin of each RNA is indicated at the top, the molecular mass of standard markers in kilobases (kb) is shown on the left.

[0032] Fig.5A: Reverse transcriptase-polymerase chain reaction (RT-PCR) analysis of EDG8 in different human endothelial cell lines (HUVECS: human umbilical vein endothelial cells; HCAEC: human coronary artery endothelial cells; HMVEC-L: human microvascular endothelial cells from lung; HPAEC: human pulmonary artery endothelial cells). EDG8-specific transcripts were detected in all endothelial cell lines. Agarose gel electrophoresis of the PCR products after 35 cycles of amplification is shown. Amplification with EDG8-specific primers yields a 522 bp EDG8-fragment as indicated by the arrow.

[0033] Fig.5B: PCR analysis of EDG8 primers for specificity of amplification of EDG8 sequences. Primers, specific for the EDG8 sequence, were checked for potential amplification of the related EDG1-7 sequences. The EDG8 specific 522 bp band occurred only when EDG8 was used as a template.

Table 1:

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	Mapping of EDG receptor	on of EDG-receptors 1-8. ors to genomic sequences allowed to derive calization of EDG1-3 was obtained from the	e a chromosomal assignment for EDG4, 5, 6 and e genecards datacollection,
10	EDG	Chromosomal localisation	according BAC AccNr.:
	EDG1	1p21.1-21.3	AL161741
	EDG2	9q31.1-32/ /18p11.3	AL157881//AP000882
	EDG3	9q22.1-q22.2	
5	EDG4	19p12	NT_000939
	EDG5	19	AC011511
	EDG6	19p13.3	AC011547
20	EDG7	1p22.3-31.2	AL139822
	FDG8	19	AC011461

#### Examples

25 Example 1: Molecular cloning of the human EDG8 receptor.

[0034] As the putative human EDG8 sequence is intronless, we cloned the receptor from human genomic DNA (CLONTECH, Palo Alto, CA, 94303-4230) via polymerase chain reaction (PCR), PCR conditions, established to amplify the EDG8 sequence were 94°C, 1 min followed by 35 cycles of 94°C, 30sec, 68C, 3 min, using GC-Melt Kit (CLON-TECH, Palo Alto, CA). Primers designed to amplify the EDG8 sequence contained a Hindlll site in the forward, and a EcoRI site in the reverse primer, respectively. The 1197 bp PCR product was cloned into the pCDNA3.1(+) mammalian expression vector (Invitrogen, Carlsbad, California) and sequenced in both directions.

Example 2: Cell culture and Transfection.

[0035] CHO-K1 cells were grown in basal ISCOVE medium supplemented with 10% fetal bovine serum at 37°C in a humidified 5% CO2 incubator. For transfections, 2 x 105 cells were seeded into 35-mm dishes. About 24 hr later cells were transiently transfected at 50-80% confluency with the indicated receptor and G-protein constructs (1µg of plasmid DNA each) using the Lipofectamine transfection reagent and the supplied protocol (GIBCO). 18-24 hr after transfection cells were seeded into 96well plates at a density of 50.000 cells per well and cultured for 18-24 additional hr until used in the functional FLIPR assays.

The cDNA for Gα16 was cloned from TF1 cells by RT-PCR and ligated into the pCDNA1.1 mammalian expression vector (Invitrogen). Murine wild type Gαq was cloned from cells by RT-PCR and inserted into the BamHI-Nsil-sites of pCDNA1.1. To create the C-terminally modified  $G\alpha_{oi5}$  subunit, in which the last five aa of wt  $G\alpha q$  were replaced with the correspoding  $G\alpha_i$  sequence, a 175-bp Bg/II-Nsil fragment was replaced, in a two piece ligation, with a synthetic DNA fragment, containing the desired codon changes. The correctness of all PCR-derived sequences was verified by sequencing in both directions.

Example 3: Fluorometric Imaging Plate Reader (FLIPR) Assay.

[0036] Twenty-four hours after transfection, cells were splitted into 96-well, black-wall microplates (Corning) at a density of 50,000 cells per well. 18-24 hr later, cells were loaded with 95ul of HBSS containing 20 mM Hepes, 2.5 mM probenecid, 4 μM fluorescent calcium indicator dye Fluo4 (Molecular Probes) and 1% fetal bovine serum for 1 h(37°C, 5% CO<sub>2</sub>). Cells were washed three times with HBSS containing 20 mM Hepes and 2.5 mM probenecid in a cell washer. After the final wash, the solution was aspirated to a residual volume of 100 µl per 96 well. Lipid ligands were dissolved in DMSO as 2 mM stock solutions (treated with ultrasound when necessary) and diluted at least 1:100 into HBSS containing 20 mM HEPES, 2.5 mM probenecid and 0.4 mg/ml fatty acid free bovine serum albumine. Lipids were

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aliquoted as 2X solutions into a 96 well plate prior to the assay. The fluorometric imaging plate reader (FLIPR, Molecular Devices) was programmed to transfer 100  $\mu$ l from each well of the ligand microplate to each well of the cellplate and to record fluorescence during 3 min in 1 second intervals during the first minute and 3 second intervals during the last two minutes. Total fluorescence counts from the 18-s to 37-s time points are used to determine agonist activity. The instrument software normalizes the fluorescent reading to give equivalent initial readings at time zero.

Example 4: Northern Blot analysis.

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[0037] Human multiple tissue Northern blots were purchased from CLONTECH (Palo Alto, CA, 94303-4230, USA) antisense RNA probes were generated by subcloning nucleotides 279-1197 of the coding region into the Bam HI-Eco RI sites of the expression vector PSPT18 (Roche Diagnostics, Mannheim, Germany) and subsequent random priming with a DIG-RNA Labeling kit (Roche Diagnostics, Mannheim, Germany), using T7 RNA polymerase. Hybridization was carried out at 68°C for 16 h in hybridization buffer (Dig Easy Hyb Roche Diagnostics, Mannheim, Germany). Each blot was washed , blocked and detected as indicated in the standard protocol with the DIG Wash and Block Buffer set (Roche Diagnostics, Mannheim, Germany) and treated with 1 ml CSPD ready-to-use(Roche Diagnostics, Mannheim, Germany) for 15 min, 37°C and developed for 5 min on the Lumiimager (Roche). Finally, each blot was stripped (50 % formamid,5% SDS, 50 mM Tris/HCl pH 7,5; 80° C, 2x 1 hour) and rehybridized with a GAPDH antisense RNA probe as an internal standard.

20 Example 5: RNA Extraction and RT-PCR.

[0038] RNA was prepared from different endothelial cell lines (HUVECS, HCAEC, HMVEC-L, HPAEC) using the TRIzol reagent (Hersteller, Lok.). Briefly, for each endothelial cell line, cells of a subconfluent 25 cm2 tissue culture flask were collected in 2,5ml TRIzol and total RNAs were extracted according to the supplied protocol. The purity of the RNA preparation was checked by veryfying the absence of genomic DNA. An aliquot of RNA, corresponding to  $\sim 5\mu g$ , was used for the cDNA generation using MMLV reverse transcriptase and the RT-PCR kit from STRATAGENE. RT-PCR was carried out in a volume of 50  $\mu$ l, the RT-PCR conditions were set to 65°C for 5 min, 15min at RT, 1 hour at 37°C, 5 min at 90°C, chill on ice.

The cDNA templates for the PCR reactions (35 cycles of 94°C for 30 sec, 68°C for 3 min) were the reverse transcribed products of RNAs isolated from human endothelial cell lines (HUVECS, HCAEC, HMVEC-L, HPAEC). Typically, 1-5 μI of reverse transcribed cDNAs were used as templates for the PCR reactions.

Example 6: Sources of materials.

[0039] 1-oleoyl-LPA, sphingosin 1-phosphate (S1P), dihydrosphingosin 1-phosphate (DHS1P), lysophosphatidyl-choline (LPC), sphingosylphosphorylcholine (SPC) and fatty acid free BSA were from SIGMA (P.O.Box 14508, St. Louis, Missouri 63178). CHO-K1 cells were obtained from the American Type culture collection (ATCC, Manassas, Virginia), cell culture media and sera from GIBCO BRL (Gaithersburg, MD), the Ca fluorescent dye FLUO4 and pluronic acid from Molecular devices (Sunnyvale CA 94089-1136,USA) human northern blot membrane from CLONTECH (1020 East Meadow Circle, Palo Alto, California 94303-4230, USA.), commercially available cDNAs (heart, fetal heart, left atrium, left ventricle, kidney, brain, liver, lung, aorta) from Invitrogen, oligonucleotides from MWG-Biotech AG (Ebersberg, Germany), the RT-PCR kit from SIGMA, the GC-melt PCR kit from Clontech (Palo Alto, CA), the expression plasmid pcDNA3.1 for EDG8 and pCDNA1.1 for expression of G-protein α subunits from Invitrogen (Carlsbad, CA 92008), competent DH5α from GIBCO and MC 1063 from Invitrogen.

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List of non-standard abbreviations:

[0041] S1P, sphingosine 1-phosphate; LPA, lysophosphatidic acid; dHS1P, dihydro sphingosine 1-phosphate; SPC, sphingosylphosphorylcholine; LPC, lysophosphatidylcholine; GPCR, G-protein-coupled receptor; G-protein, guanine nucleotide-binding protein; [Ca<sup>2+</sup>]<sub>i</sub>, intracellular Calcium concentration, RT-PCR, reverse transcription polymerase chain reaction; bp, base pair; ORF, open reading frame; EST, expressed sequence tag; FAF-BSA, fatty acid free bovine serum albumine; HUVECS. Human umbilical vein endothelial cells; HCAEC, human coronary artery endothelial cells; HMVEC-L, human microvascular endothelial cells from lung; HPAEC, human pulmonary artery endothelial cells.

Table 2:

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## SEQ ID NO. 1: Nucleotide sequence of human EDG8

15 1 ATGGAGTCGGGGCTGCGGCCGGCGCGCGGTGAGCGAGGTCATCGTCCTGCATTACAAC 61 TACACCGCAAGCTCCGCGGTGCGCGCTACCAGCCGGGTGCCGGCCTGCGCGCCGACGCC 121 GTGGTGTGCCTGGCGGTGTGCGCCTTCATCGTGCTAGAGAATCTAGCCGTGTTGTTGGTG 20 181 CTCGGACGCCACCCGCGCTTCCACGCTCCCATGTTCCTGCTCCTGGGCAGCCTCACGTTG 241 TCGGATCTGCTGGCAGGCGCCGCCTACGCCGCCAACATCCTACTGTCGGGGCCGCTCACG 25 361 GCGTCCGTGCTGAGCCTCCTGGCCATCGCGCTGGAGCGCAGCCTCACCATGGCGCGCAGG 421 GGGCCGCGCCCGTCTCCAGTCGGGGGCGCACGCTGGCGATGGCAGCCGCGGCCTGGGGC 30 541 GCTTGCTCCACTGTCTTGCCGCTCTACGCCAAGGCCTACGTGCTCTTCTGCGTGCTCGCC 35 601 TTCGTGGCCATCCTGGCCGCTATCTGTGCACTCTACGCGCGCATCTACTGCCAGGTACGC 661 GCCAACGCGCGCCCTGCCGGCACGGCCCGGGACTGCGGGGACCACCTCGACCCGGGCG 40 721 CGTCGCAAGCCGCGCTCGCTGGCCTTGCTGCGCACGCTCAGCGTGGTGCTCCTGGCCTTT 781 GTGGCATGTTGGGGCCCCCTCTTCCTGCTGCTGTTGCTCGACGTGGCGTGCCCGGCGCGC 841 ACCTGTCCTGTACTCCTGCAGGCCGATCCCTTCCTGGGACTGGCCATGGCCAACTCACTT 45 901 CTGAACCCCATCATCTACACGCTCACCAACCGCGACCTGCGCCACGCGCTCCTGCGCCTG 961 GTCTGCTGCGGACGCCACTCCTGCGGCAGAGACCCGAGTGGCTCCCAGCAGTCGGCGAGC 50 1021 GCGGCTGAGGCTTCCGGGGGCCTGCCCGCTGCCTGCCCCGGGCCTTGATGGGAGCTTC 1081 AGCGGCTCGGAGCGCTCATCGCCCCAGCGCGACGGGCTGGACACCAGCGGCTCCACAGGC 1141 AGCCCCGGTGCACCCACAGCCGCCCGGACTCTGGTATCAGAACCGGCTGCAGACTGA 55

Table 3: SEQ ID NO. 2: Amino acid sequence of human EDG8

Ε R S Ε 10 Н М F L 15 Α Α N Ι S Y R Α 20 R G R T L М С L Р Α L G N 25 K Y G G 30 T L L R G L 35 40 С C Н S С G R D S Α S G G L R R С L P G D Ρ S S Q D G D T s 45 Ŧ A Α R Т V S

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### Annex to the application documents - subsequently filed sequences linsting

[0042]

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10	Gly	Ala	Gly 35	Leu	Arg	Ala	Asp	Ala 40	Val	Val	Cys	Leu	Ala 45	Val	Суз	Ala
15	Phe	Ile 50	Val	Leu	Glu	Asn	Leu 55	Ala	Val	Leu	Leu	Val 60	Leu	Gly	Arg	His
	Pro 65	Arg	Phe	His	Ala	Pro 70	Met	Phe	Leu	Leu	Leu 75	Gly	Ser	Leu	Thr	Leu 80
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25	Gly	Pro	Leu	Thr 100	Leu	Lys	Leu	Ser	Pro 105	Ala	Leu	Trp	Phe	Ala 110	Arg	Glu
	Gly	Gly	Val 115	Phe	Val	Ala	Leu	Thr 120	Ala	Ser	Val	Leu	Ser 125	Leu	Leu	Ala
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	Val	Ser	Leu	Leu	Leu 165	Gly	Leu	Leu	Pro	Ala 170	Leu	Gly	Trp	Asn	Cys 175	Leu
40	Gly	Arg	Leu	Asp 180	Ala	Cys	Ser	Thr	Val 185	Leu	Pro	Leu	Tyr	Ala 190	Lys	Ala
45	Tyr	Val	Leu 195	Phe	Cys	Val	Leu	Ala 200	Phe	Val	Gly	Ile	Leu 205	Ala	Ala	Ile
	Суз	Ala 210	Leu	Tyr	Ala	Arg	Ile 215	Tyr	Cys	Gln	Val	Arg 220	Ala	Asn	Ala	Arg
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	Leu	Leu	Ala	Phe 260	Val	Ala	Суз	Trp	Gly 265	Pro	Leu	Phe	Leu	Leu 270	Leu	Leu
5	Leu	Asp	Val 275	Ala	Cys	Pro	Ala	Arg 280	Thr	Cys	Pro	Val	Leu 285	Leu	Gln	Ala
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15	Ile 305	Tyr	Thr	Leu	Thr	Asn 310	Arg	Asp	Leu	Arg	His 315	Ala	Leu	Leu	Arg	Leu 320
	Val	Cys	Cys	Gly	Arg 325	His	Ser	Cys	Gly	Arg 330	Asp	Pro	Ser	Gly	Ser 335	Gln
20	Gln	Ser	Ala	Ser 340	Ala	Ala	Glu	Ala	Ser 345	Gly	Gly	Leu	Arg	Arg 350	Cys	Leu
25	Pro	Pro	Gly 355	Leu	Asp	Gly	Ser	Phe 360	Ser	Gly	Ser	Glu	Arg 365	Ser	Ser	Pro
30	Gln	Arg 370	Asp	Gly	Leu	Asp	Thr 375	Ser	Gly	Ser	Thr	Gly 380	Ser	Pro	Gly	Ala
30	Pro 385	Thr	Ala	Ala	Arg	Thr 390	Leu	Val	Ser	Glu	Pro 395	Ala	Ala	Asp		
35																

## Claims

- An isolated polynucleotide comprising a nucleotide sequence that has at least 80 % identity to a nucleotide sequence encoding the polypeptide of SEQ ID NO. 2 or the corresponding fragment thereof; or a nucleotide sequence complementary to said nucleotide sequence.
  - 2. The polynucleotide of claim 1 which is DNA or RNA.
- 45 3. The polynucleotide of claim 1 wherein said nucleotide sequence is at least 80 % identical to that contained in SEQ ID NO. 1.
  - 4. The polynucleotide of claim 3 wherein said nucleotide sequence is contained in SEQ ID NO. 1.
- 50 5. The polynucleotide with sequence SEQ ID NO. 1.
  - **6.** The polynucleotide of claim 1 wherein said encoding nucleotide sequence encodes the polypeptide of SEQ ID NO. 2 or a fragment thereof.
- 7. EDG8 DNA or RNA molecule comprising an expression system wherein said expression system is capable of producing a polypeptide or a fragment thereof having at least 80 % identity with a nucleotide sequence encoding the polypeptide of SEQ ID NO. 2 or said fragment when said expression system is present in a compatible host cell.

- 8. A host cell comprising the expression system of claim 7.
- A process for producing a EDG8 polypeptide or fragment comprising culturing a host claim 8 and under conditions sufficient for the production of said polypeptide or fragment.
- 10. The process of claim 9 wherein said polypeptide or fragment is expressed at the surface of said cell.
- 11. Cells produced by the process of claim 10.

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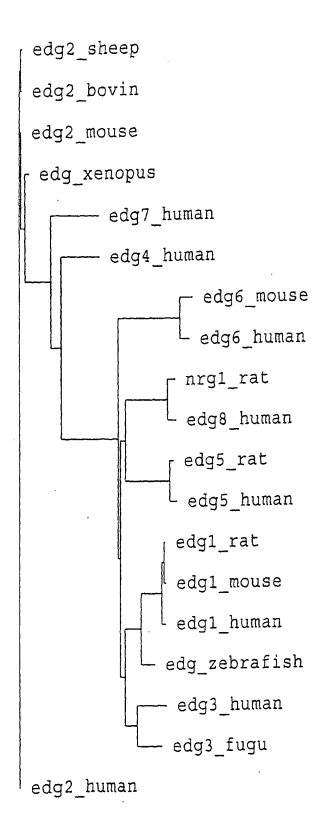
- 10 12. The process of claim 9 which further includes recovering the polypeptide or fragment from the culture.
  - 13. A process for producing a cell which produces a EDG8 polypeptide or a fragment thereof comprising transforming or transfecting a host cell with the expression system of claim 7 such that the host cell, under appropriate culture conditions, produces a EDG8 polypeptide or fragment.
  - 14. EDG8 polypeptide or a fragment thereof comprising an amino acid sequence which is at least 80 % identical to the amino acid sequence contained in SEQ ID NO. 2.
  - 15. Polypeptide of claim 14 which comprises the amino acid sequence of SEQ ID NO. 2, or a fragment thereof.
  - 16. EDG8 Polypeptide or fragment prepared by the method of claim 12.
  - 17. A process for diagnosing a disease or a susceptibility to a disease related to expression or acitivity of EDG8 polypeptide comprising:
    - c) determining the presence or absence of mutation in the nucleotide sequence encoding said EDG8 polypeptide in the genome of said subject; and/or
    - d) analyzing for the presence or amount of the EDG8 polypeptide expression in a sample derived from said subject.
  - 18. A method for identifying compounds which bind to EDG8 polypeptide comprising:
    - c) contacting a cell as claimed in claim 11 or a part thereof with a candidate compound; and
    - d) assessing the ability of said candidate compound to bind to said cells.
  - 19. The method of claim 18 which further includes determining whether the candidate compound effects a signal generated by activation of the EDG8 polypeptide at the surface of the cell, wherein a candidate compound which effects production of said signal is identified as an agonist.
  - 20. The method of claim 18 which further includes determining whether the candidate compound effects a signal generated by activation of the EDG8 polypeptide at the surface of the cell, wherein a candidate compound which effects production of said signal is identified as an antagonist.
    - 21. An agonist identified by the method of claim 19.
    - 22. An antagonist identified by the method of claim 20.
    - 23. The method of claim 18 which further includes contacting said cell with a known agonist for said EDG8 polypeptide; and determining whether the signal generated by said agonist is diminished in the presence of said candidate compound, wherein a candidate compound which effects a diminution in said signal is identified as an antagonist for said EDG8 polypeptide.
    - 24. A method as claimed in claim 23, wherein the known agonist is S1p, LPA and/or DHS1P.
- 55 **25.** An antagonist identified by the method of claim 23 or 24.
  - 26. Method of preparing a pharmaceutical composition comprising

- a) identifying a compound which is an agonist or an antagonist of EDG8,b) preparing the compound, andc) optionally mixing the compound with suitable additives.
- **27.** Pharmaceutical compound prepared by a process of claim 26.

# FIG 1A:

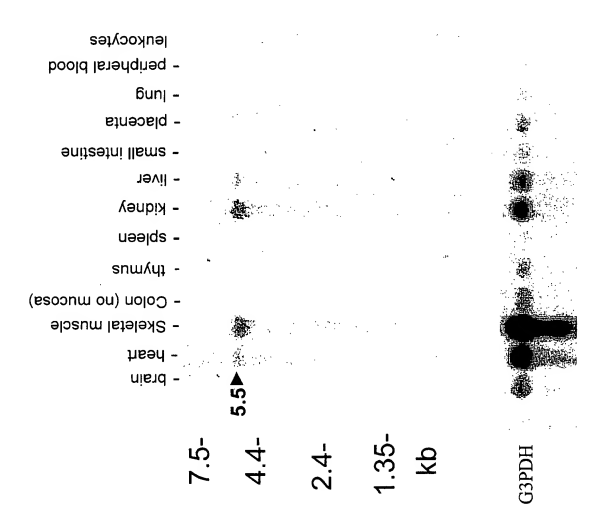
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61	TΑ	CAC	CGG	CAA	GCT	CCG	CGG	TGC	GCG	CTA	CCA	GCC	GGG	TGC	CGG	ССТ	GCG	CGC	CGA	.CGC
01	Y	Т	G	K															D	
121	GT.	CCTO	CTC	יררייי	GGC	CGT	GTG	CGC	ىسى	רבחי	CGT	יככיי	AGA	aan	ጥርጥ	AGC	CGT	GTT	'С <b>Т</b> Т	GGT
121				L																V
181	СТ	CGG	ACG	CCA	ccc	GCG	CTT	CCF	CGC	TCC	CAT	'GTT	CCT	GCT	CCT	GGG	CAG	сст	'CAC	GTT
	L	G		Н																
241	TC	GGA	гст	GCT	GGC	AGG	CGC	CGC	CTA	CGC	CGC	CAA	CAT	CCT	ACT	GTC	GGG	GCC	GCT	CACG
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•••		K		s																
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				P																
481	GT	GTC	GCT	GCT	CCT	CGG	GCT	CCI	GCC	AGC	GCI	'GGG	CTG	GAA	TTG	CCT	GGG	TCG	CCT	GGAC
	V		L																L	
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		A		W															A	
841	AC																			ACTI
	Т	-	-	V			-													
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				Ι																
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1141			CGG	TGC	ACC	CAC	AGC	CGC	ccc	GAC	TCT	GGT	ATC	AGA	ACC	GGC	TGC	AGA	CTG	Α

Fig. 1B



		1					60
	edg2 human		ISOPOFTAMN	EPOCEYNEST	AFFYNRSGKH	LAT EWNTYS	
FIG 1C:	edg7 human				DFFYNRSNTD		
	edg4 human				GFFYNNSGKE		
	edgl human				VRHYNYTGKL		
	edg3 human				REHYQYVGKL		
	edg5 human	~~~~~~~	~~~~~MGSL	YSEYLNPNKV	QEHYNYTKE.	TLETQETT	SROVASAFIV
	edg8_human	~~~~~~~	~~~~MESGL	LRPAPVSEVI	VLHYNYTGKL	RG. ARYQPGA	GLRADAVVCL
	edg6 human	~~~~MNATG	TPVAPESCQQ	LAAGGHSRLI	VLHYNHSGRL	AGR. GGPEDG	GLGALRGLSV
		61					120
	edg2_human	TVCIFIMLAN	LLVMVAIYVN	RREHEPIYYL	MANLAAADFF	AGLAYFYLMF	NTGPNTRRLT
	edg7_human				LANLAAADFF		
	edg4_human				LGNLAAADLF		
	edgl_human				IGNLALSDLL		
	edg3_human				IGNLALCDLL		
	edg5_human				LGNLAASDLL		
	edg8_human				LGSLTLSDLL		
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		121	7 D T C T T C				180
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	edg8_human						
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		301					360
	edg2 human		T.T.AFFNSAMN	DIIVEVENE	MSATFRQILC	CORSENDTER	
	edg7 human				MYGTMKKMIC		
	edg4_human						
	edgl human						
	edg3 human						
	edg5 human						
	edg8 human	PVLLQADPFL	GLAMANSLLN	PIIYTLTNRD	LRHALLRLVC	CGRHSCGRDP	SGSQQSAS
	edg6 human		ALAVLNSAVN	PIIYSFRSRE	VCRAVLSFLC	CGCLRLGMRG	<b>PGDCLARAVE</b>
		361					418
	edg2_human				~~~~~~		
	edg7_human	IPSTVLSRSD	TGSQYIEDSI	SQGAVCNKST	S		
	edg4_human						
	edgl_human				DEGDNPETIM		
	edg3_human				VKEDLPHTDP		
	edg5_human				PTFLEGNTVV		
	edg8_human						
	edg6_human	AHSGASTTDS	SLRP.RDSFR	GSRSLSFRMR	EPLSSISSVR	SI~~~~~	~~~~~

Fig. 4



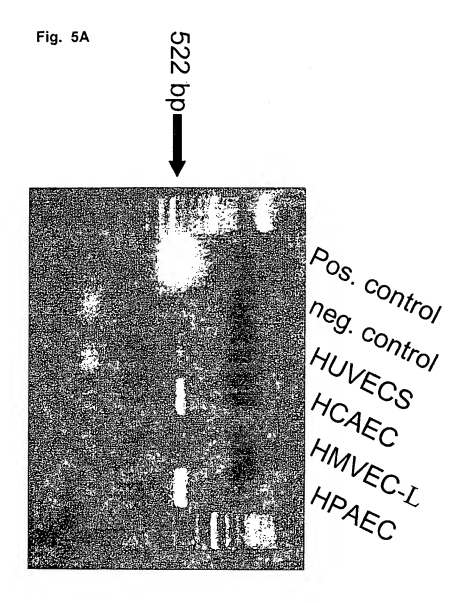
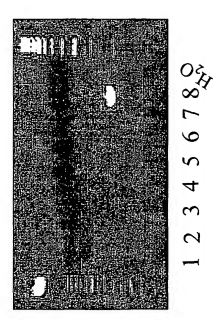


Fig. 5B





**Application Number** 

which under Rule 45 of the European Patent Convention EP  $\,00\,10\,8858\,$  shall be considered, for the purposes of subsequent proceedings, as the European search report

Category		dication, where appropriate,	Relevant	CLASSIFICATION OF THE
alegury	of relevant pass		to claim	APPLICATION (Int.Cl.7)
X	99.7 % aa seq ident SEQ ID NO:2 in 398 * the whole documen * page 15, line 13 * page 27, line 10	03-02) identity of SEQ ID :1 in 1197 bp overlap, ity of SEQ ID NO:1 with aa overlap.	1-27	C12N15/12 C07K14/705 C12Q1/68 G01N33/52 G01N33/50
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				C07K C12Q
				GOIN
INCO	MPLETE SEARCH		<del></del>	
not compt be carned	ly with the EPC to such an extent that a out, or can only be carried out partial!	application, or one or more of its claims, does a meaningful search into the state of the art or y, for these claims		
Cialmis se	arched completely			
Clams se	arched incompletely	•		
Claims no	ot searched			
Reason to	or the imitation of the search			
see	sheet C			
	Place of search	Date of completion of the search		Examiner
	THE HAGUE	13 September 2000	van	de Kamp, M
	ATEGORY OF CITED DOCUMENTS	T theory or principle E earlier patent doct	ament, but publi	
	icularly relevant if taken alone icularly relevant if combined with anoth		the application	
	ument of the same category	L i document cited for	r other reseance	



# INCOMPLETE SEARCH SHEET C

Application Number EP 00 10 8858

Reason for the limitation of the search:

Present claims 21, 22, 25 and 27 relate to agonists and antagonists for EDG8. Support within the meaning of Article 84 EPC and/or disclosure within the meaning of Article 83 EPC is to be found, however, for only a small proportion of the compounds claimed. In the present case, the claims so lack support, and the application so lacks disclosure, that a meaningful search over the whole of the claimed scope is impossible. Consequently, in view of the description, the search has been carried out for those compounds which appear to be supported and disclosed by the description, namely sphingosine-1-phosphate (SIP), lysophosphatidic acid (LPA), dihydrosphingosine-1-phosphate (DHS1P), sphingosylphosphorylcholine (SPC), lysophosphatidylcholine (LPC), and G-protein alpha-subunits.



Application Number

EP 00 10 8858

	DOCUMENTS CONSIDERED TO BE RELEVANT	CLASSIFICATION OF THE APPLICATION (Int.CI.7)	
Category	Citation of document with indication, where appropriate. of relevant passages	Relevant to claim	
	DATABASE EMBL 'Online!  EBI; ID ACO11461, AC ACO11461, 8 October 1999 (1999-10-08) "Homo sapiens chromosome 19 clone CTC-429L19, WORKING DRAFT SEQUENCE, 4 unordered pieces" XP002147034 Note: 100.0 % nt seq identity of of nt 90281-91477 with SEQ ID NO:1 in 1197 bp overlap, 100.0 % aa seq identity of translated nt 90281-91477 with SEQ ID NO:2 in 398 aa overlap. * the whole document *	1-6	
X,D	GLICKMAN M ET AL.: "Molecular cloning, tissue-specific expression, and chromosomal localization of a novel nerve growth factor-regulated G-protein-coupled receptor, nrg-1."  MOL. CELL. NEUROSCI., vol. 14, no. 2, August 1999 (1999-08), pages 141-152, XP000939383  Note: 80.1 % nt seq identity wit. EQ ID NO:1 in 1046 bp overlap, 85.8 % aa seq identity with SEQ ID NO:2 in 401 aa overlap.  * abstract *  * page 142, right-hand column, line 43 - page 144, left-hand column, line 20 *	1-3, 6-11, 13-25	TECHNICAL FIELDS SEARCHED (Int.Cl.7)

23

EPO FORM 1503 03.82 (P04C10)



Application Number

EP 00 10 8858

	DOCUMENTS CONSIDERED TO BE RELEVANT		CLASSIFICATION OF THE APPLICATION (Int.CI.7)
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	
X	WO 00 22129 A (ARENA PHARMACEUTICALS INC; LIAW CHEN W (US); BEHAN DOMINIC P (US);) 20 April 2000 (2000-04-20) Note: 99.7 % nt seq identity of SEQ ID NO:31 with SEQ ID NO:1 in 1197 bp overlap, 99.7 % aa seq identity of SEQ ID NO:32 with SEQ ID NO:2 in 398 aa overlap.  * page 32, line 9-21 *  * table D *	1-20,26	
Α	WO 99 19513 A (LXR BIOTECHNOLOGY INC ;ERIKSON JAMES (US); KIEFER MICHAEL (US); GO) 22 April 1999 (1999-04-22) * page 29, line 1 - page 33, line 15 *	1-27	TECHNICAL FIELDS SEARCHED (Int.Cl 7)
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